PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference	·						
M/45016-PCT	FOR FURTHER ACTION	See Form PCT/IPEA/416					
International application No. PCT/EP2005/001038	International filing date (day/month/year) 02.02.2005	Priority date (day/month/year) 03.02.2004					
International Patent Classification (IPC) or r INV. A61K31/366 A61K31/40 A61K	national classification and IPC (31/22 A61K31/695 A61K9/20 A61P9	9/10					
Applicant FERRER INTERNACIONAL, S.A.	et al.						
This report is the international pre- Authority under Article 35 and tra	eliminary examination report, established insmitted to the applicant according to Ar	d by this International Preliminary Examining rticle 36.					
2. This REPORT consists of a total	2. This REPORT consists of a total of 5 sheets, including this cover sheet.						
3. This report is also accompanied by ANNEXES, comprising:							
a. 🛛 sent to the applicant and t	to the International Bureau) a total of 3	sheets, as follows:					
	ion, claims and/or drawings which have ting rectifications authorized by this Autho	been amended and are the basis of this report ority (see Rule 70.16 and Section 607 of the					
☐ sheets which supersed beyond the disclosure Supplemental Box.	de earlier sheets, but which this Authorit in the international application as filed, a	ty considers contain an amendment that goes as indicated in item 4 of Box No. I and the					
sequence listing and/or tab	Bureau only) a total of (indicate type and loles related thereto, in electronic form on ing (see Section 802 of the Administrative	number of electronic carrier(s)), containing anly, as indicated in the Supplemental Box re Instructions).					
4. This report contains indications re	elating to the following items:						
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☑ Box No. I Basis of the report☐ Box No. II Priority	ort						
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	ent of opinion with regard to novelty, invention	entive step and industrial applicability					
 □ Box No. IV Lack of unity of invention □ Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement 							
☐ Box No. VI Certain docume	_						
☐ Box No. VII Certain defects i	in the international application						
	tions on the international application						
	Date of completion	n of this report					
Date of submission of the demand		·					
Date of submission of the demand 30.11.2005	19.06.2006						
	19.06.2006						

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/EP2005/001038

B	ox No. I	Basis of the repo	rt ·
1. W	/ith regar	d to the language, th	nis report is based on
\boxtimes	the int	ternational application	n in the language in which it was filed
	a trans	slation of the internat anslation furnished fo	ional application into , which is the language or the purposes of:
	☐ inte	ernational search (un blication of the interna	der Rules 12.3(a) and 23.1(b)) ational application (under Rule 12.4(a)) v examination (under Rules 55.2(a) and/or 55.3(a))
Ně	ave been	rurnisnea to the rece	If the international application, this report is based on (replacement sheets which eiving Office in response to an invitation under Article 14 are referred to in this re not annexed to this report):
De	escription	Page	
1-1	•	, rages	as originally filed
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	aims, Nur	mbers	
24			as originally filed
1-2	23		received on 30.11.2005 with letter of 30.11.2005
Dra	awings, S	Sheets	
1/1			as originally filed
	a sequ	ence listing and/or an	y related table(s) - see Supplemental Box Relating to Sequence Listing
3. 🗆	The am	nendments have resu	Ilted in the cancellation of:
		description, pages	
		claims, Nos. drawings, sheets/figs	
	☐ the	sequence listing (spe	ecify):
	□ any	table(s) related to se	quence listing (specify):
4. □ had Sup	a not bee	oort has been establis n made, since they h al Box (Rule 70.2(c))	shed as if (some of) the amendments annexed to this report and listed below ave been considered to go beyond the disclosure as filed, as indicated in the .
		description, pages	•
		claims, Nos. drawings, sheets/figs	
	☐ the s	sequence listing (spe	
	⊔ any t	table(s) related to sed	quence listing (specify):
*	If ite	m 4 applies, so	me or all of these sheets may be marked "superseded."

Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

3: 1

Novelty (N)

Yes: Claims

1-23

No: Claims

Inventive step (IS)

Yes: Claims

No: Claims 1-23

Industrial applicability (IA)

Yes: Claims

1-23

No: Claims

2. Citations and explanations (Rule 70.7):

see separate sheet

10/588377 IAP11 Rec'd PCT/PTO 02 AUG 2006 International application No.

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

PCT/EP2005/001038

Re Item V

Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- 1. The documents cited in the International Search Report (ISR) are consecutively numbered D1-D8 in the order of their listing. If not indicated otherwise, reference is made to the passages cited in said ISR.
- 2. Document D1 relates to pharmaceutical compositions comprising a statin. Said formulations may optionally contain an anti-foaming agent such as simethicon in an amount of about 0% to about 0.5% of the final formulation (D1, p.30, paragraph [075]). However, the only example which illustrates the addition of simethicone is example 9, which describes a pH-independent coating which comprises approximately 0.15% by weight of simethicone emulsion as dispersant. Having regard to the fact that said proportion refers to the coating only (and not to the entire formulation comprising pravasatin) and further refers to an emulsion of simethicone (and not simethicone per se), the actual proportion of simethicone in dosage forms having the coating of example 9 will be far below 0.15% by weight. Taking into account that the amount of pravasatin in said formulation is well above 5% by weight (see examples 1-5), the weight ratio of simethicone versus pravastatin is expected to be below 0.25.

Document D2 describes the composition of commercial film-coated tablets comprising atorvastatin. Simethicone emulsion is one of a number of inactive ingredients contained in said tablets. Simethicone has the function of an auxiliary agent and therefore the weight ratio of simethicone versus atorvastatin is believed to be far below 0.25.

Thus, the subject-matter of claims 1-23 is not novel in the sense of Art. 33(2) PCT.

3. The problem to be solved by the present invention may be regarded as the provision of an improved hypocholesterolemic composition comprising a statin which does not cause flatulence.

The solution to the problem posed is the addition of an antiflatulent agent (simethicone or dimethicone) in a weight ratio of antiflatulent agent versus statin of at least 0.25 in order to achieve an antiflatulent effect.

It is well-known that simethicone has antiflatulent effects and that it can be added to compositions comprising other active ingredients (see D4, D5). D1 explicitly points to the optional addition to compositions comprising pravastatin of anti-foaming agents such as simethicone in an amount of about 0% to about 0.5% of the final formulation (p.30, first paragraph).

The applicant states that the proportion of simethicone in D1 is too low to achieve an antiflatulent effect. However, in the absence of any evidence showing an unexpected effect in relation to the claimed weight ratio an inventive step cannot be acknowledged, Art. 33(3) PCT.

10/588377

CLAIMS

IAP11 Rec'd PCT/PTO 02 AUG 2006

- 1. A pharmaceutical composition comprising a statin and an antiflatulent agent wherein the weight ratio of antiflatulent agent versus statin is at least 0.25.
- 2. The composition of claim 1 wherein the ratio is at least 1.50.
- The composition of any one of claims 1 and 2 wherein the statin is selected from the group consisting of atorvastatin, cerivastatin, fluvastatin, lovastatin, pitavastatin, pravastatin, rosuvastatin and simvastatin, or pharmaceutically acceptable salts and hydrates thereof.
- 4. The composition of claim 3 wherein the statin is simvastatin or a pharmaceutically acceptable salt thereof.
- 5. The composition of any one of claims 1 to 4 wherein the antiflatulent agent is selected from the group consisting of simethicone and dimethicone.
- 6. The composition of claim 4 wherein the antiflatulent agent is simethicone.
- 7. The composition of any one of claims 1 to 6 wherein the composition is a tablet, capsule, syrup, solution, powder, granule, or emulsion.
- 8. The composition of claim 7 wherein the tablet is a coated tablet.

- 9. The composition of claim 8 wherein the coated tablet comprises a core and a coating, the core comprising the statin and the antiflatulent agent.
- 10. The composition of any one of claims 7 and 8 wherein simvastatin is present in an amount from 2.5 to 100 mg per tablet.
- 11. The composition of claim 10 wherein simvastatin is present in an amount from 5 to 80 mg per tablet.
- 12. The composition of any one of claims 7 and 8 wherein simethicone is present in an amount from 25 to 250 mg per tablet.
- 13. The composition of claim 12 wherein simethicone is present in an amount of 125 mg per tablet.
- 14. The composition of any one of claims 1 to 13 further comprising one or more diluents, one or more binders, one or more disintegrants and one or more lubricants.
- 15. The composition of claim 14 wherein the diluent is selected from the group consisting of microcrystalline cellulose and their derivatives, lactose, mannitol, calcium phosphates, starch, and the mixtures thereof.
- 16. The composition of claim 14 wherein the binder is selected from the group consisting of starch, polyethylene glycols, polyvinylpyrrolidones, cellulose derivatives, and the mixtures thereof.
- 17. The composition of claim 14 wherein the disintegrant is selected from the group consisting of colloidal

silicon dioxide, croscarmellose, polyvinylpyrrolidone, starch and its pregelatinized derivatives, and the mixtures thereof.

- 18. The composition of claim 14 wherein the lubricant is selected from the group consisting of talc, magnesium stearate, stearic acid, sodium stearyl fumarate, PEG 8000, and the mixtures thereof.
- 19. The composition of any one of claims 1 to 18 further comprising one or more antioxidants and one or more wetting agents.
- 20. The composition of claim 8 wherein the coating of the tablet comprises a cellulose derivative or its pharmaceutically acceptable salt, an acrylic polymer, triethyl citrate, titanium dioxide and one or more lubricants.
- 21. The composition of claim 17 wherein the cellulose derivative is hydroxypropyl methylcellulose.
- 22. The composition of any one of claims 1 to 21 further comprising one or more colouring agents.
- 23. A process for preparing a composition according to any one of claims 1 to 22 by direct compression of components thereof.